FROM : McGREGOR&ADLER, P. C.

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REMARKS

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Oath/Declaration

The oath or declaration has been objected to as defective because the Combined Declaration and Power of Attorney submitted February 17, 1998 does not identify that the instant application is a continuation-in-part of 08/702,205. A replacement Combined Declaration and Power of Attorney has been submitted herewith listing the application numbers of the applications to which the instant application claims priority. The Applicant respectfully requests that the objection to the oath or declaration be withdrawn.

The 35 U.S.C. §103(a) Rejection

Claims 8-15 remain rejected under 35 U.S.C. §103(a) as being unpatentable over Tanimoto et al. (Leukemia, 3:339-348, 1989) or Scheinberg et al. (Leukemia, 3:440-445, 1989) in view of Thorpe et al. (Immunological Revie0ws, 62:119-158, 1982), Andrews et al. (Blood, 62:124-132, 1983) and Rosenblum et al. (U.S. Patent No. 5,631,348, Filed August 14, 1990). These rejections are respectfully traversed.

Tanimoto et al. and Scheinberg et al. teach an anti-CD33 monoclonal antibody M195. Thorpe et al. teach conjugation of antibodies to gelonin. Andrews et al. teach anti-CD33 antibodies may be useful for the treatment of leukemia. Rosent lum et al. teach the sequence of gelonin. Nevertheless, Applicant respectfully submits that combining these references does not lead to the claimed invention, nor do they encompass all the important features of the present invention.

Although the references teach M195 and gelionin, they do not specifically teach M195 conjugated to recombinant gelonin.

Tanimoto et al., Scheinberg et al. (Leukemia, 3:440-445, 1989),

Thorpe et al. and Andrews et al. do not teach or suggest the making and using of M195 conjugated to recombinant gelonin, whereas Rosenblum et al. do not teach or suggest conjugating recombinant gelonin to M195 for cancer treatment. While the above combination of references might suggest that it would be obvious to try to conjugate M195 to gelonin to form an effective immunotoxin, one skilled in the art would have no method of knowing beforehand a functional immunotoxin would result from

FROM : McGREGOR&ADLER, P. C. Available CARYNE NO. :

such a conjugation. It is possible that the conjugation event could either alter the binding site of the antibody or reduce the toxicity of the gelonin conjugate. Further time-consuming, pon-routine experimentation would be necessary to show that a functional immunotoxin was produced.

In addition, for immunotoxins to be effective, they must be effectively internalized into the target cell, and they must be directed to the correct intracellular compartment in order for the toxin to kill the cell. For example, uptake of the toxin into the lysosomal compartments may lead to the enzymatic digestion of the toxin before it can have any effect on the cell. Because the actual cytotoxicity of antibody-cytotoxin conjugates vary as 4 result of complex factors, it would not have been obvious beforehand that a functional immunotoxin would result from the conjugation of M195 to gelonin without additional, undue experimentation. In contrast. the present invention has detailed disclosure on the effectiveness of the M195-recombinant gelonin conjugate in vitro (Examples 9, 10) and in vivo (Example 14). Applicant also has evidence that indicate efficacy and lack of significant toxicity of the M195-recombinant gelonin conjugate in clinical setting.

FROM : McGREGOR&ADLER, P. Be vailable Copyone NO. :

The Applicant respectfully submits that no teaching, suggestion or incentive may be gleaned from the references, individually or together, with regard to the making and using of the claimed invention. Therefore, the invention as a whole is not prima facie obvious to one of ordinary skill in the art at the time the invention was made. Accordingly, Applicant respectfully requests that the rejection of claims 8-15 under 35 U.S.C. §103 (a) as obvious over Tanimoto or Scheinberg (Leukemia, 3:440-45, 1989) in view of Thorpe, Andrews and Rosenblum be withdrawn.

Claims 8-15 remain rejected under 35 U.S.C. §103(a) as being unpatentable over Scheinberg (U.S. patent 5,730,982, March 24, 1998) in view of Thorpe et al. (Immunological Reviews, 62:119-158, 1982), Andrews et al. (Blood, 62:124-132 1983) and Rosenblum et al. (U.S. Patent No. 5,631,348, Filed August 14, 1990). This rejection is respectfully traversed.

Thorpe et al., Andrews et al. and Rosenblum et al. have been discussed in detail above. U.S. patent 5,730,982, issued to Scheinberg, teaches the use of M195-radioisotopes conjugates

FROM : McGREGOR&ADLER, P. C. Vailable Copy NO. :

However, the cited patent does not teach, in cancer treatment. suggest or show data on the making and using of the M195recombinant gelonin conjugate for cancer treatment. As with the combination of Thorpe et al., Andrews et al. and Rosenblum et al. with Tanimoto or Scheinberg (Leukemia, 3:440-445, 1989), the combination of Thorpe et al., Andrews et al. and Rosenblum et al. with Scheinberg (U.S. patent 5,730,982) provides evidence that M195 conjugated to gelonin will result in an effective immunotoxin for cancer treatment. Once again, it is possible that the conjugation event could either alter the binding site of the antibody or reduce the toxicity of the gelonin conjugate or that problems may arise in the effective internalization conjugation into the target cells. While the combined references may suggest that it is obvious to try to form an effective immunotoxin by conjugating M195 to gelonin, additional, undue experimentation would be necessary to show that the formed immunotoxin was effective. Accordingly, Applicant respectfully requests that the rejection of claims 8-15 under 35 U.S.C. §103(a) as obvious over the combination of Scheinberg (U.S. patent 5,730,982) Thorpe, Andrews and Rosenblum be withdrawn.

FROM : McGREGOR&ADLER, P. C. Be vailable Copy No. :

This is intended to be a complete response to the Office Action mailed July 13, 2000. If any issues remain outstainding, the Examiner is respectfully requested to telephone the uindersigned attorney of record for immediate resolution.

Respectfully submitted,

Date: Nov 1, 2000

Benjamin Aaron Adler, Ph.D., J.D.

Registration No. 35,423 Counsel for Applicant

McGREGOR & ADLER, LLP 8011 Candle Lane Houston, Texas 77071 (713) 777-2321 badler1@houston.rr.com